Mass spectrometric quantification of Candida albicans surface proteins to identify new diagnostic markers and targets for vaccine development

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Chapter 6:

Surface stress induces a conserved cell wall stress response in the pathogenic fungus *Candida albicans*

Introduction

During infection, *C. albicans* encounters stresses from host defenses (*e.g.* fever, oxidative and nitrosative stress), environmental niches (*e.g.* hypoxia in the gut, antimicrobial peptides in saliva and in epithelial layers) and antifungal intervention (*e.g.* azoles and echinocandins). Many of these stresses directly affect the cell surface, which has distinct fungal features compared to mammalian cells. The most important difference is the presence of a cell wall.

The cell wall is the initial site of host-pathogen interaction and composed of a skeletal layer of carbohydrates, mainly β-glucans and chitin, which is covered with an external layer of covalently anchored mannoproteins. These mannoproteins have been shown to serve a variety of functions, from immune-evasion (22, 41) and nutrient acquisition (3, 77) to adhesion, biofilm formation (28, 72), and tissue degradation (65). Many cell wall proteins also directly modulate the wall composition and architecture as carbohydrate-active enzymes (14). Reinforcement of the cell wall in response to antifungal stresses is well described, especially an increase in chitin content as a result of increased chitin synthesis (49, 76). The cell wall proteome itself is highly dynamic (35) and adaptable in response to external conditions (68, 70) as well as morphological changes (26). This dynamic surface is crucial for an opportunistic pathogen, enabling it to colonize different niches in a variety of hosts. Sites of infection differ dramatically in, for example, oxygen levels, pH, and available nutrients. Another environmental factor that has a major impact on the fungal surface and growth is temperature, which can vary considerably depending on the host species. While most fungi are not able to grow above 40°C (62), *C. albicans* causes infections in many animals (29) among them birds (*e.g.* penguins, pigeons), whose body temperature is in this range (60, 66). As a consequence of prolonged thermal stress both *S. cerevisiae* as well as *C. albicans* cells have been shown to accumulate trehalose, which facilitates proper protein folding under stress conditions (4, 44). In *S. cerevisiae* thermal stress also leads to the activation of the cell wall integrity (CWI) pathway which in turn affects the composition of the wall and its proteins (reviewed in (39)). In addition, Mkc1, the *C. albicans* ortholog of Slt2 in *S. cerevisiae*, and a key signal transducer in the CWI pathway of *C. albicans*, has been suggested to be required for growth at elevated temperatures (52). Another member of the CWI pathway, Pkc1, was shown to be involved in the response to fluconazole (37), a widely used antifungal agent that leads to the depletion of ergosterol in the fungal membrane (50), resulting in increased membrane fluidity (1, 12).
We recently established that fluconazole does not only elicit membrane stress, but also leads to a stressed cell wall. Since membrane fluidity is correlated with temperature (79), we hypothesized that a similar effect as seen for fluconazole could be achieved by growth at elevated temperatures. In this study, we examined the effect of thermal stress on the wall composition, the secretome, and the cell wall proteome as well as underlying regulatory proteins. Relative quantification revealed a set of wall remodeling proteins that was induced during thermal stress. Intriguingly, the same set of proteins was shown previously to be involved in the response to other surface stressors (7, 68), suggesting a general stress response. Therefore, we investigated the phosphorylation status of three stress-responsive MAP kinases, Hog1, Cek1, and Mkc1. This analysis revealed that strongly increased Mkc1 phosphorylation is a hallmark of prolonged thermal stress as well as long-term exposure to wall and membrane stress. We also show that high-temperature stress as well as antifungal treatment leads to cell clustering, which is linked to reduced secretion of chitinases, since the clusters are dispersible by exogenous chitinase, but not by SDS treatment or vortexing. To cope with prolonged surface stress a highly conserved wall remodeling response, probably mainly mediated by Mkc1 signaling is activated and the wall is reinforced with chitin through increased chitin synthesis and reduced chitin degradation. The identification of proteins involved in wall stress adaptation opens the door to the development of strategies to improve clinical outcomes of antifungal treatment.

Prolonged high temperature stress causes cell wall stress

*C. albicans* is able to withstand and grow in conditions many other fungi cannot survive. This suggests that *C. albicans* has evolved strategies to minimize the impact of these stresses. We previously showed that the widely used antifungal fluconazole leads to both membrane as well as wall stress, which the cell tries to counteract by increased chitin incorporation as well as the increase of repair-associated proteins in the wall (68). Similar to fluconazole, temperature affects the membrane composition and fluidity, which could in turn affect the polysaccharide composition of the cell wall and the incorporation levels of its covalently anchored proteins. When analyzing the effects of temperature on growth at two pHs, we found that liquid cultures grew faster at pH 4 than at pH 7.4 at all temperatures tested and that at both pHs growth was peaking at 30°C (Table 6.2). Compared to 30°C, growth at 42°C was reduced by ~70% at pH 4 and >80% at pH 7.4. Interestingly, when grown on solid medium at 42°C and pH 4,
the cells were considerably more resistant to this temperature than at pH 7.4 (Figure 6.1).

<table>
<thead>
<tr>
<th>Growth temp.</th>
<th>Biomass (mg dry weight/ml)</th>
<th>pH 7.4</th>
<th>pH 4</th>
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<tbody>
<tr>
<td>25°C</td>
<td>1.7 ± 0.2</td>
<td>2.2 ± 0.3</td>
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</tr>
<tr>
<td>30°C</td>
<td>4.4 ± 0.3</td>
<td>5.0 ± 0.4</td>
<td></td>
</tr>
<tr>
<td>37°C</td>
<td>2.0 ± 0.1</td>
<td>3.7 ± 0.1</td>
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<tr>
<td>42°C</td>
<td>0.7 ± 0.1</td>
<td>1.4 ± 0.5</td>
<td></td>
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</tbody>
</table>

Table 6.2. Biomass yields after 18 h at pH 7.4 and pH 4

1Mean ± SD (n = 3)

Figure 6.1. Spot assays of Candida albicans grown at 37°C and 42°C in the presence and absence of chemical stressors.

1:10 serial dilutions of SC5314 overnight cultures were spotted onto YNB-S plates at pH 4 or pH 7.4 and supplemented with 100 mg/L SDS, 50 mg/L Calcofluor white or 1 mg/L Congo red, respectively, and grown for 2 days.

It is known that cells respond to wall stress by increasing the chitin levels in the wall, thus strengthening the weakened cell wall (38). We therefore analyzed the chitin content of walls from cells grown at 37°C and 42°C, both at pH 7.4 and pH 4. At pH 7.4 chitin levels increased about 1.7-fold and at pH 4 the amount of chitin in the cell wall even doubled at 42°C compared to the 37°C control (Table 6.3). Overall, growth at pH 4 led to higher wall chitin levels compared to pH
7.4 (Table 6.3), which probably contributes to the higher heat resistance at low pH and might be related to morphology (Figure 6.1). Furthermore, when *C. albicans* was incubated at 42°C in the presence of the wall-perturbing compounds Calcofluor white and Congo red or the membrane perturbant SDS (Figure 6.1), higher sensitivity was observed, supporting the notion that cell wall integrity is compromised at elevated temperatures.

<table>
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<tr>
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<th>Average ± SD¹</th>
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<tr>
<td>pH 4 37°C</td>
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</tr>
<tr>
<td>pH 4 42°C</td>
<td>97.4 ± 8.0**</td>
<td></td>
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<tr>
<td>pH 7.4 37°C</td>
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<tr>
<td>pH 7.4 42°C</td>
<td>54.8 ± 9.0**</td>
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</table>

Table 6.3. Cell wall chitin content after growth for 18 h.

¹Mean ± SD (n ≥ 3) ** p < 0.01 vs. 37°C

Prolonged high temperature stress affects the wall proteome

Subsequently, we analyzed the cell wall proteome at 42°C by applying relative quantification using a metabolically labeled ¹⁵N-reference culture and FTMS (26, 68). Combining technical and biological variance, the median coefficient of variation of our measurements was 11.2% and 13.1% for pH 4 and pH 7.4, respectively. Comparing the wall proteomes obtained at 37°C and 42°C revealed a strong impact of prolonged heat treatment at pH 7.4, while the changes were less pronounced at pH 4 (Figure 6.2). At both pHs the wall levels of Sap9, involved in maintaining wall integrity (2), increased. The wall levels of the chitin transglycosylases Crh11 and Utr2 (9) and of a protein of unknown function involved in wall integrity, Ecm33 (42), also increased significantly at both pHs. The levels of other GPI-anchored wall proteins clearly decreased at both pHs upon heat stress, namely, the chitinase Cht2, the adhesin Als2, and the small temperature-regulated protein Rhd3. Interestingly, the set of proteins that showed reduced wall levels upon thermal stress also included the GPI-anchored wall proteins Csa1, Pga10, and Rbt5. They contain one or more CFEM domains, consisting of eight cysteines separated by conserved spacing (36) and have been linked to iron starvation and heme/hemoglobin binding (77, 78). Fascinatingly, both of the antagonistically pH-regulated transglycosylases Phr1 and Phr2 are strongly increased in abundance at pH 7.4, suggesting a relief of the
Rim101-regulated repression of \textit{PHR2} at this pH during wall stress (13, 70). In summary, our observations reveal that upon prolonged thermal stress the wall proteome changes dramatically, showing significantly increased levels of wall proteins involved in wall maintenance and transglycosylation. This same set has been also identified during fluconazole stress (68), hinting at a more general stress response.

**Figure 6.2. Relative quantification of wall proteins using FTMS and \textsuperscript{15}N-labeling.**

The graphs are representing log\textsubscript{2} ratios of wall proteins at 42°C versus 37°C, at both pH 4 (graph on the left, median CV\%=13.1\%) and pH 7.4 (graph on the right, median CV\%=11.2\%). Positive values represent an increase and negative values a decrease of the corresponding protein at 42°C compared to the 37°C control.
Figure 6.3. Immunoblot analysis of phosphorylated MAP kinases. The cultures were grown for 18 h at pH 7.4 and 37°C in the presence of various chemical stressors (0.5 mg/L fluconazole, 125 mg/L SDS, 5 mg/L Congo red, and 40 mg/L Calcofluor) or subjected to thermal stress at 42°C. Fifteen µg of protein extracts were loaded per lane. Pgk1 was used as a loading control. The anti-phospho-p38 MAPK (Thr180/Tyr182) antibody or anti-phospho-p44/42 MAPK (Thr202/Tyr204) antibody was used to detect Hog1~P and Mkc1~P/Cek1~P, respectively.

Thermal stress leads to Mkc1 phosphorylation

Next, we wanted to investigate the involvement of the major MAP kinase pathways in *C. albicans* in the late response to continuous thermal stress in comparison to the late responses elicited by known wall- and membrane-perturbing compounds. We therefore carried out immunoblot analysis of the phosphorylated forms of Mkc1, Cek1, and Hog1 after 18 h of growth under the tested conditions. As shown in Figure 6.3 the phosphorylated form of Mkc1 became more abundant after prolonged thermal stress and even more so in the presence of fluconazole, Congo red, and Calcofluor white. This is consistent with earlier observations that loss of *MKC1* results in a strongly decreased resistance to Congo red and Calcofluor white (51). The levels of Cek1~P tended to increase only slightly, whereas Hog1~P did not seem to increase significantly at 42°C and decreased in the presence of the other stress conditions. Under our growth conditions Mkc1 seems to play an important role in the late responses to membrane and wall stress while Cek1 and Hog1 play only minor roles. Our observations do however not exclude a role for these MAP kinases during earlier responses to thermal stress (44).
**Figure 6.4.** Temperature, stress conditions and chitinase treatment influence the morphology of *C. albicans* wild type and mutant strains.

*C. albicans* wild type and mutants were grown in YNB-S at pH 7.4 and were stained with Calcofluor white. All cultures were grown at 37°C except when labeled otherwise. + denotes treatment with 0.5 U chitinase for 3 h. FCZ: Fluconazole (0.5 mg/L); CFW: Calcofluor white (50 mg/L); CR: Congo red (2 mg/L); SDS: Sodium Dodecyl Sulfate (100 mg/L). The scale bar indicates 10 µm.

Stress-induced cell clustering is linked to decreased chitinase abundance

Fluorescence microscopy in combination with Calcofluor white staining revealed significant formation of cell clusters at 42°C compared to lower growth temperatures (Figure 6.4). These cell clusters could not be disrupted by SDS treatment or sonication, suggesting a covalent linkage between the cells. The same holds for chitinase knockout strains and Δ/Δ ace2 and Δ/Δ cbk1 mutants, which are involved in the regulation of cell separation (data not shown). This is also consistent with the intense staining by the chitin-binding dye Calcofluor white of the septal region (Figure 6.4). Mass spectrometric analysis of the secretome from cells grown at pH 4 revealed a significant decrease of the cell separation-related secretory proteins Cht3 and Eng1 at 42°C compared to lower temperatures (Table 6.4; for all >40 detected proteins see supplemental tables S3
Due to the large number of intracellular proteins in the secretome of cells grown at 42°C in pH 7.4 medium these data were not used for comparison (data not shown). The presence of large number of intracellular proteins is most likely due to increased breakage or lysis of stressed cells and hyphae, hinting at a higher resilience of pH 4-grown yeast cells. Interestingly, as already mentioned above, the GPI-anchored, wall-bound chitinase Cht2 also decreased upon thermal stress especially at pH 4 (Figure 6.2).

<table>
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<tr>
<th>Name</th>
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</tr>
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<tbody>
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<td></td>
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<td>30°C</td>
</tr>
<tr>
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</tr>
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</tr>
<tr>
<td>Xog1</td>
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<td>7.3</td>
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Table 6.4. Semiquantitative analysis of selected medium proteins
The values represent the average relative abundances (%) of protein identifications of selected medium proteins per growth condition as estimated by peptide counting from Q-TOF analysis. For information on all >40 identified proteins see supplemental tables S3 and S4. n.d., not detected.

Cht3 and Eng1 - both less abundant during heat stress - are involved in the dissolution of wall carbohydrates after the formation of a primary septum, allowing cell separation. Cht2 and Cht3 are both chitinases, close homologs to the only chitinase in *Saccharomyces cerevisiae*, Cts1, with Cht3 being the major chitinase in *Candida albicans* (18). Eng1 is an endo-1,3-beta-glucanase and also involved in cell separation (20). Knockout mutants of *cht3* and the homozygous diploid mutant strain *cht2/cht3* showed cellular non-separation even at 37°C (Figure 6.5). The protein kinase Cbk1 and the transcription factor Ace2, which participate in the RAM pathway (64), are major regulators of wall and secreted proteins. The expression of *CHT3* and *ENGL* both decrease about threefold
when $ACE2$ is knocked out (48). The complete absence of cell separation in the $ace2$ deletant strain (34) suggests that Cht3 and Eng1 act synergistically during cell separation (Figure 6.4). Like the knockouts or cells exposed to thermal stress, cells treated with the membrane-perturbing compounds fluconazole and SDS as well as with the cell wall-perturbing agents Calcofluor white and Congo red displayed a similar non-separation phenotype (Figure 6.4), suggesting that this phenotype is associated with plasma membrane and cell wall stress. To corroborate that decreased chitin degradation is the reason for the cell separation defect as observed during cell wall stress, we treated $C. albicans$ with exogenous chitinase for 3 h. As shown in Figure 6.4, treatment with chitinase led to the dispersal of cell clusters, most notably for Δ/Δace2 and Δ/Δcbk1.

Figure 6.5. Relative dispersion of different $C. albicans$ mutant strains and wild type cells cultured under various stress conditions.

The sedimentation time of wild type cells grown at 37°C without (Wt 37°C) or with 0.5 mg/L fluconazole (FCZ) or 40 mg/L Calcofluor white (CFW), of wild type cells grown at 42°C, or of various mutant strains at 37°C was compared. Strains that were treated with chitinase are marked with +. The time needed for the optical density to decline to 80% of the initial optical density was determined and expressed as percentage of wild type cells grown at 37°C (relative dispersion). Bars represent averages ± standard deviation of two experiments; * $p < 0.05$, ** $p < 0.01$ relative to wild type control at 37°C.

For a comparative analysis of cell clustering we introduced a spectrophotometric assay to measure relative sedimentation times. For each culture, the time required for the OD$_{600}$ to decline to 80% of its initial value was measured and their sedimentation times were expressed as percentage of the sedimentation
time of wild type cells grown at 37°C. As expected, Δ/Δace2 and Δ/Δcbk1 cells, which cluster strongly, sedimented much faster than wild type cells grown at 37°C (Figure 6.5). Cultures grown in the presence of Calcofluor white or fluconazole also showed smaller relative sedimentation times, as did the Δ/Δcht3 and Δ/Δcht2/3 homozygous diploid mutant strain, but not the Δ/Δcht2 strain, underlining the importance of Cht3 for cell separation and the dispersion of cell clusters. Importantly, treatment of the Δ/Δace2 and Δ/Δcbk1 strains with exogenous chitinase lead to strongly increased sedimentation times. Cells grown at 42°C also sedimted significantly faster than control cells and this phenotype could be largely restored by treating them with exogenous chitinase. This supports the notion that cell clustering during membrane and cell wall stress is linked to the decrease in chitinase levels in the medium.

Discussion

In this study we have shown that prolonged thermal stress is causing wall stress, which activates the Mkc1 signaling pathway. Chitin levels of the cell wall become higher leading to reinforcement of the wall, whereas chitinase levels in the medium decrease. This probably reflects decreased formation and secretion of Cht3 leading to incomplete degradation of the primary chitinous septum during cytokinesis, thus leading to reduced cell separation. Additionally, a set of wall maintenance and repair proteins is strongly increased on the wall. All these responses seem to be a general consequence of membrane and wall stress. As previously demonstrated for the antifungal fluconazole, we could validate that prolonged thermal stress is causing wall stress. Both fluconazole and high temperatures are affecting membrane fluidity (58, 79), which probably indirectly weakens the cell wall. Cells with a wall defect are less resistant to the chitin-binding dye Calcofluor white, the β-1,3-glucan-binding Congo red, and SDS (53). Since C. albicans grown at 42°C showed less resistance to these compounds, this supports that thermal stress causes wall stress. Another hallmark of a stressed cell wall is an increase in wall chitin levels. This wall reinforcement in response to various stresses is a well described phenomenon in fungi (11, 21, 56, 57, 76). While only a minor constituent of the cell wall, an increase of a few percent of chitin leads to a more robust wall. For example, in the presence of the β-1,3-glucan inhibitor caspofungin, the resulting loss of structural cell wall integrity is compensated by an increase of chitin in the wall (76). An increase of chitin can be achieved by either increased expres-
sion of the chitin synthases (CHS), a decrease in the degradation of chitin by the chitinases (CHT) or, most likely, both. A variety of pathways are involved in cell wall maintenance including the Mk c1, Hog1, Msb2-Cek1 and calcineurin pathways as well as the chaperone Hsp90, which all have been shown to be involved in the regulation of chitin synthase expression (33, 37, 49).

Less attention has been focused on the degradation of chitin and its role in stress resistance. Cells grown at 42°C for 18 h showed a severe cell separation defect. Similarly, cell aggregates were also described for several mutants that show a defective wall, like the glycosylation mutants mmn9, pmt1 and pmt4, the wall protein mutants ecm33, pga10 and pga13 and mutants in the wall-associated transglycosylase bgl2 (23, 42, 54, 59, 71). Gregori et al. assigned the cell separation defect in caspofungin-treated cells to an increased expression of adhesins, mainly Als1 (25). We also observed an increase of Als1 in stressed cells, but found that the lack of cell separation is not primarily due to protein-protein interactions, since the cell clusters could not be dispersed by SDS and vortexing or by mild sonication. In addition, our mass spectrometric analysis did reveal a distinct decrease of Cht3 and Eng1 in the secretome (Table 6.4) and cell clusters could be dispersed by adding exogenous chitinase (Figure 6.4).

For full cell separation of mother and daughter cell the β-glucan layer needs to be severed and the primary septum dissolved. The observed reduced levels of chitinases and endoglucanase are presumably not sufficient anymore to ensure proper cell segregation, thus resulting in cell clustering. We hypothesized that generally wall and membrane stress lead to an increase of chitin by a concerted increase of chitin synthesis as well as reduced degradation. We therefore also tested SDS and Calcofluor White and observed the same clustering phenotype (Figure 6.4). In conjunction with our previous analyses of fluconazole- and Congo red-treated cells (68), this suggests that the decreased abundance of cell separation enzymes, the increase of chitin in the wall and the absence of cell separation in stressed cells are directly linked and represent a general stress response. Furthermore, we observed a similar clustering phenotype when testing knockout mutants of the chitinase Cht3, endoglucanase Eng1, and severe clustering of their regulators, Ace2 and Cbk1 (Figure 6.5). Cht3 has been described as the major chitinase in C. albicans (18) and since Ace2 regulates both ENG1 and CHT3 (48), the more severe phenotype of the Ace2 knockout mutant suggests a synergistic effect between Cht3, Eng1, and other potential Ace2 targets. Strikingly, an ace2 mutant is less sensitive to Calcofluor white, SDS, and azoles (27). Possibly, within cell aggregates a microenvironment can develop that of-
fers protection against these compounds and this might be further facilitated by
the formation of extracellular matrix material (30). The growth rates of ace2
mutant cells and wild type cells at 42°C are however similar (27).
In a recent study, in which C. albicans was treated with micafungin, as expected
an increased expression of the chitin synthases was observed, but in addition a
marked down-regulation of CHT2 and CHT3 was reported (31). Resistance to
echinocandins is mainly mediated by the mutation of FKS1, the targeted subunit
of β-1,3-glucan synthase (55). Intriguingly, a recent analysis of echinocandin-
resistant strains without FKS1 mutations also identified mutations in both CHT2
and CHT3, with two point mutations in CHT3 directly located in the active site
and the substrate binding cleft, conceivably severely limiting the activity of the
enzyme (17). These strains also exhibited a strongly increased chitin content,
which has been shown to be leading to a higher resistance to caspofungin in vivo
(38). This suggests that mutations that render chitinases less active lead to
increased resistance to wall stress. Taken together, we propose that different
forms of wall and membrane stress result in diminished cell separation. Not just
the increased expression of chitin synthases, but also the concerted reduction of
chitin degradation by reduced abundance of chitinases results in wall reinforce-
ment with chitin and diminished cell separation. This is supported by the disper-
sal of cell clusters in the presence of chitinase and the associated slower sedi-
m entation.
Increased chitin incorporation is only one of a number of stress responses at the
disposal of C. albicans. At any given time, carbohydrate-active enzymes are
remodeling the wall structure and architecture (14). Among these are the afore-
mentioned chitinases (Cht2, Cht3), the calcineurin-dependent chitin transglyco-
sylases (Crh11, Utr2) (53), β-glucan transglycosylases (Pga4, Phr1, Phr2) as
well as the β-1,3-glucan crosslinking Pir1 and the yapsin-like protease Sap9.
Intriguingly, while the chitinases are reduced in abundance in response to stress,
all others are strongly increased in abundance during thermal stress (Figure 6.2).
We also observed a marked increase of these proteins in response to fluconazole
and Congo red stress (68). The yapsin-like protease Sap9 has been implicated to
have a major role in cell wall integrity and host pathogen interaction (2). Con-
sistent with our data presented here and previously (68), SAP9 expression has
been reported to be strongly increased in response to antifungal treatment (10).
Pir1 has been shown to be crucial for cell wall maintenance (43), while the chi-
tin transglycosylases Crh11 and Utr2, both belonging to the same protein fami-
ly, are required for cell wall biogenesis. Although Phr1 and Phr2 have been re-
ported to be differently expressed according to pH (47), our data show that both proteins are strongly increased at pH 7.4 during stress conditions consistent with our previous results (68), suggesting a relief of Rim101 control of *PHR2* at neutral pH. Taken together and including the results of our previous work (68) and transcriptional data obtained for cells treated with a variety of antifungal agents (7, 40), we hypothesize that this set of proteins participates in a highly conserved response to wall and membrane stress.

Secreted aspartyl proteases, most notably *SaP8* (61, 63, 74), have also been implicated in processing Msb2, a transmembrane protein which upon cleavage activates the Cek1 MAP kinase pathway. A recent study also showed that proteolytic cleavage of Msb2 is not only important for signaling, but that its extracellular part can directly interact with human antimicrobial peptides (73), hinting at an important role in host-induced stresses. We observed strong Mkc1 phosphorylation under the stress conditions tested. Mkc1 and its homolog Slt2 in *S. cerevisiae* and *C. glabrata* have been shown to be involved in heat resistance (52), virulence (16), wall reinforcement (11) and caspofungin resistance (80). We show here that Mkc1 is not only transiently activated by wall and membrane stress, but that Mkc1 continues to be activated in the presence of stress. This suggests that *C. albicans* adapts to long-term cell surface stress with a continuous activation of Mkc1, resulting in increased expression of remedial proteins, including the previously mentioned carbohydrate-active enzymes.

Concluding remarks

In our previous work (68, 69) we identified a core set of abundantly secreted proteins (*Cht3, Mp65, Scw11, Sim1, Sun41, Tos1, Xog1*) under virtually all growth conditions tested. When grown at 25°C, 30°C and 37°C both at pH 7.4 or pH 4 and at pH 4 at 42°C most of these proteins were secreted in abundance as well (Table 6.3). Nevertheless *Cht3* was considerably less abundant at pH 4 and 42°C and *Scw11* at pH 4 and 25°C. Since this core set of proteins seems to be consistently present and ample peptide identifications suggest abundance, these proteins or antibodies raised against them might be suitable potential infection markers. The increase of repair-associated proteins during heat stress (Figure 6.2) has also been observed during clinically induced stresses like fluconazole (68) or caspofungin treatment (7). They seem to be a general and relevant response to membrane and wall stress, which *C. albicans* also encounters
in the host in the form of various antifungal peptides, like histatins, alpha- and beta-defensins and cathelicidins. We have shown that thermal stress causes wall stress in \textit{C. albicans}, supported by the increased chitin levels and sensitivity to wall perturbing agents at 42°C. Furthermore we have shown that the wall stress response to thermal treatment seems to be similar to various surface stressors, leading to the activation of \textit{Mkc1} signaling, an increase of certain repair-associated wall proteins, reduction in cell-separation enzymes, most importantly \textit{Cht3}, leading to wall reinforcement with chitin and a correlated decrease in cell separation.
Material and Methods

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<td>SC5314</td>
<td>ace2Δ::FRT/ace2Δ::FRT</td>
<td>ΔΔ ace2</td>
<td>(34)</td>
</tr>
<tr>
<td>CAMM-292-4</td>
<td>CAI4</td>
<td>$ura3/ura3\Delta::hisG/cbk1Δ::hisG$</td>
<td>ΔΔ cbk1</td>
<td>(45)</td>
</tr>
</tbody>
</table>

Table 6.1. Strains used in this study.

Strains, growth conditions, biomass measurements, and microscopy

All chemicals were obtained from Sigma-Aldrich unless otherwise stated. *C. albicans* wild type SC5314 or mutant strains (Table 6.1) were pre-cultured overnight at 30°C in liquid YPD medium (10 g/L yeast extract, 20 g/L peptone and 20 g/L glucose) in a rotary shaker at 200 rpm. The next day, flasks containing 50 ml YNB-S (6.7 g/L Yeast Nitrogen Base (YNB), 20 g/L sucrose) either buffered at pH 7.4 using 75 mM MOPSO [3-(N-morpholino)-2-hydroxypropanesulphonic acid] or at pH 4 using 75 mM tartaric acid were inoculated from the overnight cultures to an OD$_{600}$= 0.05. Sucrose was used as a carbon source to avoid glucose repression. MOPSO and tartaric acid were selected as buffers because they are hydrophilic and non-metabolizable with pKₐ values close to the desired pH (69). The cultures were stirred at 200 rpm and incubated for 18 h at either 25°C, 30°C, 37°C or 42°C. For analyzing the response to different wall or membrane stresses the culture media were supplemented with Calcofluor white (CFW), Congo red (CR), fluconazole (FCZ) or sodium dodecyl sulfate (SDS). To increase biomass yield for wall purification of 42°C-grown cells, multiple flasks were pooled per biological replicate. To determine biomass of the cultures after 18 h of growth, cultures were spun down and the pellets were dried at 60°C overnight and weighed the next day. For morphological analysis, cells were stained with CFW and photographed using fluorescence microscopy. For staining, 3 µl of the culture were mixed on a glass slide with 3 µl of a 1:1,000 dilution of a 10 mg/ml stock of CFW, incubated for 5 min, and visualized using an Axiovert 40 CFL microscope (Zeiss, UK). To determine the effect of exogenous chitinase on various strains, the cells were incubated for 3 h at room tempera-
ture in the presence of 0.5 U chitinase purified from *Trichoderma viride* (C8241) in 50 mM phosphate buffer (pH 6.1) and then stained and visualized as described above.

**Spectrometric assay of relative sedimentation times**

*C. albicans* cells were cultured as described above. *C. albicans* wild type SC5314 or mutant strains (Table 6.1) were grown for 18 h at 37°C or 42°C, in the presence or absence of fluconazole or Congo red. Subsequently, strains indicated with + in Figure 6.5 were treated with chitinase as described above. The OD_{600} from 800 µl of a culture with or without chitinase treatment was measured each min for 15 min. Each strain and growth condition was measured in duplicate and the time required for the optical density to decline to 80% of the original value, designated as the sedimentation time, was determined for each time course. The sedimentation time of wild type cells grown at 37°C was set at 100%, and the sedimentation times of all other conditions and strains were expressed as % of this value.

**Spot assays to test resistance to cell wall-perturbing agents**

*C. albicans* wild type or mutant strains were grown overnight in liquid YPD medium at 200 rpm and 30°C. The overnight culture was 1/10 serially diluted and five µl of undiluted culture and each consecutive ten-fold diluted culture were spotted on plates containing YNB-S at pH 7.4 solidified with 1.5% agar. For treatment with cell wall- or plasma membrane-perturbing agents the plates were supplemented with CFW, CR or SDS. After incubation at 37°C or 42°C for two days the plates were photographed.

**Intracellular protein extraction and immunoblot analysis**

*C. albicans* was cultured in YNB-S at 37° or 42°C as described above. After 18 h the cells were harvested by centrifugation at 4°C and washed once with cold PBS. Intracellular proteins were prepared as described previously (46). Briefly, cells were resuspended in cold lysis buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 0.5% of the detergent NP-40, 2 mg/L leupeptin, 2 mg/L pepstatin, 1 mM PMSF, 2 mM Na_3VO_4, 50 mM NaF). Cold glass beads with a diameter of 0.25-0.50 mm were added and cells were disintegrated using the Precellys 24 homogenizer (Bertin Technologies, France). After centrifugation to remove glass beads and cell debris the protein concentration of the supernatants was determined and normalized using the Bradford assay (6). Proteins were separated on a 4-12% Bis-Tris gel (Invitrogen, San Diego, CA) and transferred to a PVDF membrane (Millipore, Billerica, MA). Membranes were blocked in a Tris-Buffered Saline-Tween solution (TBS-T) with 10% BSA and 50 mM NaF at room temperature for 30 minutes. They were further incubated overnight at 4°C with either anti-phospho-p38 MAPK (Thr^{180}/Tyr^{182}) antibody or anti-phospho-p44/42 MAPK (Thr^{202}/Tyr^{204}) antibody (both from New England Biolabs, UK) in TBS-T with 5% BSA and 50 mM NaF. As a loading control, antibodies against 3-phosphoglycerate kinase (Acris, Germany) were
used. After washing the PVDF membranes five times for five minutes in TBS-T they were incubated with peroxidase-conjugated goat-anti rabbit antisera (Thermo Fisher Scientific, Waltham, MA) in TBS-T with 5% BSA and 50 mM NaF at room temperature for 1 hour. Subsequently, the membranes were washed three times for five minutes in TBS-T. Proteins were visualized using ECL plus detection reagents (GE Healthcare, Waukesha, WI).

Secretome analysis
Secretome analysis was performed as described before (68, 69). Cells were separated from the growth medium by centrifugation. In order to remove remaining cells the supernatant was passed through a 0.2-μm filter and then concentrated using 10-kDa cut-off filters (Amicon Ultra-15 Centrifugal filter units, Millipore, Billerica, Mass.). Secreted proteins were quantified using the BCA assay and BSA as standard, and normalized to dry weight. For mass spectrometric analysis the concentrated proteins were further reduced, alkylated, and digested with trypsin as described for wall proteins.

Cell wall purification, reduction, and alkylation
Cell walls were prepared as described elsewhere (14) with some modifications. Briefly, the cell pellet was washed several times with PBS, and then subjected to breakage in a Precellys 24 homogenizer with glass beads (0.25-0.50 mm) in the presence of a protease inhibitor cocktail. Full breakage was controlled by light-microscopic inspection. The pellet was washed several times with 1 M NaCl and then washed twice with MilliQ-water. The pellets were resuspended in an appropriate volume of SDS extraction buffer (150 mM NaCl, 20 g/L SDS, 100 mM Na-EDTA, 100 mM β-mercaptoethanol, 50 mM Tris-HCl, pH 7.8) and incubated while shaking at 200 rpm at 37°C for 30 min. The pellets were spun down and resuspended in fresh SDS extraction buffer and incubated overnight under the same conditions. These steps ensure thorough and stringent washing to retain only covalently anchored wall proteins. The following day, the pellets were boiled four times for ten min in SDS extraction buffer, washed several times with MilliQ-water and lyophilized overnight.

The purified wall pellets were stored at -80°C until needed or directly reduced and S-alkylated as described elsewhere (82). Briefly, treatment of the wall pellets with reducing solution (10 mM dithiothreitol in 100 mM NH₄HCO₃) for 1 h at 55°C is followed by alkylation (65 mM iodoacetamide in 100 mM NH₄HCO₃) for 45 min at room temperature in the dark and subsequent quenching of the reaction by adding 55 mM dithiothreitol/100 mM NH₄HCO₃ for five min. To eliminate excess iodoacetamide and dithiothreitol, the pellets were washed six times with 50 mM NH₄HCO₃. Similarly, the secretome samples were reduced and alkylated on 10-kDa cut off spin filters (Amicon, Billerica, MA) (69). Reduced and alkylated samples were either directly trypsinized or stored at -80°C until used.
Determination of cell wall protein and chitin content

The wall protein content and the chitin content of the cell walls were determined as described previously (32). Four mg of freeze-dried cell walls were suspended in 100 μl 1 M NaOH and boiled for ten minutes. After cooling to room temperature the suspension was neutralized by the addition of 100 μl 1 M HCl and centrifuged to get rid of insoluble material. Using a BCA assay the protein content of the supernatant was determined and compared to bovine serum albumin as a standard.

For chitin determination the insoluble cell wall components obtained after the NaOH extraction step were hydrolyzed in one ml 6 M HCl for 17 h at 100°C. After evaporation of the liquid under a stream of nitrogen the pellet was resuspended in one ml demineralized water. To 100 μl sample 100 μl 1.5 M Na2CO3 in 4% acetylacetone were added and the mixtures were boiled for 20 minutes. Samples were cooled to room temperature and 700 μl 96% ethanol and 100 μl of 1.6 g of p-dimethyl-aminobenzaldehyde in 30 ml of concentrated HCl and 30 ml of 96% ethanol were added. After a 1-h incubation at room temperature absorbance at 520 nm was measured and compared to a standard of glucosamine.

Metabolic 15N-labeling of the reference cultures

Two different culturing conditions were used to generate a 15N-metabolically labeled reference culture. Cells grown at pH 4 and 30°C and at pH 7.4 and 37°C were labeled as described previously (26). Briefly, optimal 15N-loading was ensured by pre-culturing twice in YNB-S with 15N-labeled ammonium sulfate as the sole nitrogen source (Spectra Stable Isotopes; 15N content >99%) with a starting OD 600 of 0.05. The final pre-culture was then used to inoculate the appropriate media for the reference culture and cells were grown at 30°C or 37°C and 200 rpm for 18 h. The 15N-labeled cells of both cultures were harvested; their walls were isolated, freeze-dried, combined in a 1:1 ratio based on dry weight, divided in 2-mg aliquots and stored at -80°C.

Sample preparation for MS analysis and 14N/15N mixing

Sample preparation for MS analysis was performed as described previously (26, 68, 69). For Q-TOF LC-MS/MS analysis of secreted proteins, alkylated proteins were digested for 18 h using 2 μg Trypsin Gold (Promega, Madison, WI, USA) and subsequently desalted using a C18 tip column (Varian, Palo Alto, CA, USA) according to the supplier’s instructions. For relative quantification, 2 mg of the freeze-dried, reduced, and alkylated 14N-query walls were mixed 1:1 based on protein content with 15N-reference walls (with 60% reference walls at pH 7.4 and 80% at pH 4) and resuspended in 50 mM NH4HCO3 and then digested and desalted as described above. To eliminate acetonitrile, all samples were evaporated using a Speedvac (Genevac, Ipswich, England) and the peptides were resolubilized in 20 μl of 0.1% trifluoroacetic acid (TFA). The amount of peptides in all
samples was determined using a NanoDrop ND-1000 (Isogen Life Science, IJsselstein, The Netherlands) at 205 nm as described before (8, 15, 26, 68).

LC-MS/MS (Q-TOF) analysis and protein identification
Two hundred and fifty ng of peptides in 10 µl 0.1% TFA were injected onto an Ultimate 2000 nano-HPLC system (LC Packings, Amsterdam, The Netherlands) equipped with a PepMap100 C18 reversed phase column (75 µm inner diameter, 25 cm length; Dionex, Sunnyvale, CA). A linear gradient with increasing acetonitrile concentration (0% - 50%) over 45 min and an elution flow rate of 0.3 µl/min was used for LC separation. The eluate was on-line ionized by electrospray in a Q-TOF (Micromass, Whyttenshawe, United Kingdom). Survey scans were acquired from m/z 350-1500. For low energy collision-induced dissociation (MS/MS), the most intense ions were selected in a data-dependent mode. Pkl (peak list) files were generated using the MaxEnt3 algorithm, included in the Masslynx Proteinlynx software. Proteins were identified by submitting the pkl files to an internally licensed version of MASCOT (Matrix Science, Great Britain), searching against a complete ORF translation of the C. albicans genome. In MASCOT two miscleavages and a tolerance of 0.5 Da for peptides and MS/MS were allowed. Based on probabilistic MASCOT scoring a P value of <0.05 was considered significant for peptide identification. Four independently obtained biological samples (the biological replicates) were analyzed for each condition. Each biological sample was subjected to two MS/MS runs (technical replicates). All identified proteins were subjected to signal peptide prediction using SignalP3.0 (5) and prediction of a GPI anchor sequence using the BIG-PI fungal predictor (19). For a semi-quantitative analysis of our data, we calculated for each growth condition the % peptide identifications. Per biological replicate the number of peptide identifications per protein was divided by the total number of identified peptides. The % spectral counts were averaged per growth condition.

LC-FT-MS/MS data acquisition
MS/MS data were acquired using an ApexUltra Fourier transform ion cyclotron resonance mass spectrometer (Bruker Daltonic, Bremen, Germany) equipped with a 7T magnet and a nano-electrospray Apollo II DualSource™ coupled to an Ultimate 3000 (Dionex, Sunnyvale, CA, USA) HPLC system. Samples containing 80 ng of the 14N and 15N tryptic peptide mixtures were injected as a 3-µl 0.1% trifluoroacetic acid aqueous solution and loaded onto the PepMap100 C18 (5-µm particle size, 100-Å pore size, 300-µm inner diameter x 5-mm length) precolumn. Following injection, the peptides were eluted via an Acclaim PepMap 100 C18 (3-µm particle size, 100-Å pore size, 75-µm inner diameter x 250-mm length) analytical column (Thermo Scientific, Etten-Leur, The Netherlands) using a linear gradient from 0.1% formic acid / 6% CH3CN / 94% H2O to 0.1% formic acid / 40% CH3CN / 60% H2O over a period of 120 min at a flow rate of 300 nL/min.
Data dependent Q-selected peptide ions were fragmented in the hexapole collision cell at an Argon pressure of $6 \times 10^{-6}$ mbar (measured at the ion gauge) and the fragment ions were detected in the ICR cell at a resolution of up to 60,000. Mass calibration was better than five ppm over all MS and MS/MS spectra in the LC-MS/MS chromatogram. MS/MS rate was about 1.3 Hz. This yielded more than 9,000 MS/MS spectra over the 120-min LC-MS/MS chromatogram.

LC-FTMS/MS data processing and relative protein quantification

Raw FT-MS/MS data were processed with the MASCOT DISTILLER program, version 2.4.3.1 (64 bits), MDRO 2.4.3.0 (MATRIX science, London, UK), including the Search toolbox and the Quantification toolbox. Peak-picking for both MS and MS/MS spectra were optimized for the mass resolution of up to 60,000. Peaks were fitted to a simulated isotope distribution with a correlation threshold of 0.7, with minimum signal to noise of 2. MS/MS spectra from precursor peptides within a mass tolerance of 0.03 Dalton and within a retention time window of maximum 60 seconds were added before peak-picking.

The processed data were searched with the MASCOT server program 2.3.02 (MATRIX science, London, UK) against a complete *Candida albicans* ORF translation (6210 entries) obtained from CGD (www.candidagenome.org). Trypsin was used as enzyme and two missed cleavages were allowed. Carbamidomethylation of cysteine was used as a fixed modification and oxidation of methionine as a variable modification. Semitryptic peptides were allowed. The peptide mass tolerance was set to five ppm and the peptide fragment mass tolerance was set to 0.01 Dalton. The quantification method was set to the Metabolic $^{15}$N labeling method to enable MASCOT to identify both $^{14}$N and $^{15}$N peptides.

Using the quantification toolbox, the isotopic ratios for all identified proteins were determined as average of the isotopic ratios of the corresponding light over heavy peptides. Selected critical settings were: Method was set to average protein ratio type, requiring bold red and a significance threshold of 0.05 with the precursor protocol type selected. The $^{15}$N loading value was set to 99.4 via the components correction for elements and the report ratio was set to L/H. For integration, the Simpsons integration method with the survey integration source was used. Elution time shift was allowed with an elution time delta of 20 seconds and a standard error threshold of 0.15. The correlation threshold based on isotopic distribution fit was set to 0.98. All charge states were used with an XIC threshold of 0.1 and a maximal XIC width of 200 seconds. Only unique peptides were used for quantification and a peptide threshold of 0.05 together with the least homology threshold for peptides. Normalisation mode was set to none and outliers were automatically removed.
LC-FTMS/MS data were acquired and analyzed of four replicas of both 37°C and 42°C cell cultures at pH 4, and of three replicas of both 37°C and 42°C cell cultures at pH 7.4. For all these 14 analyses the comprehensive MASCOT DISTILLER quantification reports are published as an EXCEL documents in the supplementary information. Only bona fide wall proteins previously verified were used for quantification (Figure 6.2; Supplemental tables 1 and 2). These reports show that over all analyses, cell wall proteins are quantified as L/H isotopic ratios averaged over a maximum of 25 corresponding tryptic peptides. From the geometric standard deviations it can be estimated that errors in the protein mean average isotopic ratios are limited to less than 15%.

For Als1, Phr2, Sap9, and Tos1 the changes in the protein levels between the 37°C and 42°C culture were so large that for the two conditions different tryptic peptide sets were identified. To obtain more overlap between these tryptic peptide sets for the two conditions, the corresponding missing peptides were manually identified from the chromatogram MS trace, based on their accurate mass and LC-retention and their L/H isotopic ratios were determined. These peptide isotopic ratios were then included in the averaged isotopic ratios for the above proteins.

Tables S1 and S2 summarize the cell wall protein L/H isotopic ratios of each replica sample. The results show that the isotopic ratios over the replicas are very consistent. This indicates that the biological variation is not dominant, the cell culture and cell processing are reproducible, and the \(^{14}\text{N}/^{15}\text{N}\) cell culture mixing errors are limited. Therefore the final protein isotopic ratios \(^{14}\text{N}/^{15}\text{N}\) in tables S1 and S2 are replica averages. The log\(_2\) values of the resulting \(^{14}\text{N}/^{15}\text{N}\) protein isotopic ratios are listed in the histograms of Figure 6.2, which shows the perturbation in the cell wall proteome upon increasing the temperature from 37°C and 42°C for pH 4 and pH 7.4.

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